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Filling reference libraries with diatom environmental sequences: strengths and weaknesses

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Diatom species identification with DNA metabarcoding is an economical, fast and reliable alternative to identification via light microscopy for river quality monitoring. Using a short DNA sequence of the *rbcL* gene and 'Diat.barcode', a reference barcode library, enables the identification of more than 90% of the environmental sequences to species level in French rivers. But the completeness of this library is much lower in other regions, such as the tropical French overseas departments. A barcode library completion method using high-throughput sequencing data combined with microscopy count data from natural samples (Rimet et al. 2018) was applied and tested in rivers of Martinique and Guadeloupe (West Indies), for which only 45% of the environmental sequences could be identified to species level using Diat.barcode v9. Assigning barcodes to the most abundant species in the islands by this method is illustrated with *Ulnaria goulardii* and two new species belonging to *Nupela* and *Epithemia*, which are also described in this paper. The more complex situation of morphologically similar species is illustrated by reference to *Gomphonema designatum* and *G. bourbonense*. Using a combination of molecular and morphological data, their conspecificity, as *G. bourbonense*, is demonstrated with their reference barcodes. However, when several morphologically similar species and several environmental sequences belonging to the same clade are present, it is not possible to relate the barcodes to corresponding morphological species.

Applying this method enabled the Diat.barcode library (v.10) to be updated, with 84% of the environmental sequences from the West Indies now identifiable at the species level. However, many morphological species still lack barcodes. In these cases, more classical methods, such as cell isolation, Sanger sequencing and morphological observations of cultures, must be applied.

Keywords: diatoms, taxa, morphology, DNA, genetic, phylogeny

Introduction

Given their value as ecological indicators, diatoms are now required for the assessment of water ecosystems quality in Europe (e.g. Kelly et al. 2014) as part of the Water Framework Directive (European Commission, 2000) and in the USA under the Clean Water Act (e.g. Barbour et al. 1999, Potapova & Charles 2007, Hausmann et al. 2016). To apply diatoms as monitoring tools, species have to be identified and counted and then biotic indices calculated (Rimet 2012). These methods are standardized, and identification and counting are carried out using microscopy (CEN 2014). However, there are some difficulties to identify

species with light microscopy. It requires time and trained analysts with good knowledge of the taxonomic literature, but inter-analyst variation can often influence the final result (Kahlert et al. 2009). In addition, many semi-cryptic species and species complexes (e.g. Trobajo et al. 2009, Kermarrec et al. 2012, Abarca et al. 2014, Kelly et al. 2015, Pinseel et al. 2020) make morphological identification complex.

An alternative solution is the DNA barcoding approach (Hebert et al. 2005), in which specific short DNA sequences, so-called barcodes, enable diatom species to be identified. DNA metabarcoding (Pompanon et al. 2011),

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based on High-Throughput-Sequencing (HTS), expanded the barcoding concept to environmental samples, in which species mixtures can be identified from natural samples. Several studies have successfully applied this approach to diatoms (Kermarrec *et al.* 2013, Visco *et al.* 2015, Zimmermann *et al.* 2015, Vasselon *et al.* 2017a, Rivera *et al.* 2018a, b). To enable efficient, accurate species identification based on DNA barcodes, reliable reference barcoding libraries are required. These libraries connect DNA barcodes to species and are used to identify the species present in samples. Several curated libraries for protists exist (e.g. PR2 for microbial diversity in Guillou *et al.* [2013], Phytool for phytoplankton in Canino *et al.* [2021]) and for diatoms Diat.barcode, which is open access and has been maintained since 2012 (Rimet *et al.* 2019). Reference barcodes in Diat.barcode come from two sources: (1) the NCBI nucleotide database and (2) unpublished sequences of culture collections. The chosen barcode for this reference library is *rbcL*, a chloroplast gene marker suitable for species-level identification of diatoms (Kermarrec *et al.* 2013, 2014). The last version of the Diat.barcode reference library (v9) contained 8066 sequences from 1491 species in 300 genera. Diat.barcode is almost complete for some regions of Europe, like France, where 91% of the environmental sequences from rivers can be identified to species level (Rimet *et al.* 2021). However, this library is still largely incomplete for the French tropical region, where diatom biomonitoring also must be applied for routine river assessment. This was the case for the French West Indies (Guadeloupe and Martinique islands), where only 45% of the environmental sequences from samples from the river monitoring network in 2018 and 2019 could be identified with Diat.barcode (Rimet *et al.* 2019). There is, therefore, a need to expand this reference database if metabarcoding will be used for diatom monitoring in rivers.

There are several methods for filling diatom reference libraries (Rimet *et al.* 2018): (1) single-cell isolation and culturing, followed by Sanger sequencing (e.g. Evans *et al.* 2007, Trobajo *et al.* 2009, Abarca *et al.* 2014, Zimmermann *et al.* 2014), (2) single-cell PCR followed by microscope observation of the living or oxidized frustule for species identification (Takano & Horiguchi 2006, Gomez *et al.* 2012, Hamilton *et al.* 2015, Khan-Bureau *et al.* 2016, Lefebvre *et al.* 2017, Skibbe *et al.* 2018, Hamilton *et al.* 2019), (3) direct Sanger sequencing of environmental samples with very low diversity, such as a *Didymosphenia* M. Schmidt bloom in mountain streams (Jaramillo *et al.* 2015), (4) use of HTS data from environmental samples and comparison to light microscopy (LM) and scanning electron microscopy (SEM) analyses to identify barcodes for target species (Rimet *et al.* 2018). This last method has the advantage that HTS enables sequencing of several hundreds of samples in a single run, generating several millions of sequences at reasonable cost and with good sequencing quality (Pfeiffer *et al.* 2018). However, linking

the sequences to the diatom species observed in LM and SEM remains the main challenge.

This last approach was used in this study, with several examples from Guadeloupe and Martinique, where river samples were sequenced with Illumina MiSeq and observed with LM and SEM. The objective of the study is to show that this approach is easily applicable in some cases, even if species are new to science. However, there are more complex situations, and even some for which this approach is not applicable. We analyse these various cases and highlight the reasons for the success or failure of the approach.

Materials and methods

Study site: Martinique (1128 km²) is located in the volcanic arc of the Lesser Antilles, in the Caribbean Sea, between Dominica to the north and Saint Lucia to the south. Guadeloupe (1436 km²) is located in the Caribbean archipelago, between the Tropic of Cancer and the equator. These islands have a tropical climate with two distinct seasons, a generally dry season from December to June and a wet winter season from July to December.

Sampling: The sampling was carried out in 2018 and 2019 during the dry season. One hundred and one samples were collected from the two study areas. The samples were collected from March to June (for Martinique: 12–16 May 2018 and 25–28 March 2019 and 24 June 2019; for Guadeloupe: 15–23 June 2018 and 10–17 April 2019). The sampling procedure followed European standards (European Committee for Standardization 2014a, 2014b). Benthic diatoms were collected from at least five stones in fast-flowing parts of rivers. The upper surfaces of the stones were scrubbed with a clean toothbrush to collect biofilms. The samples were then fixed with ethanol (final concentration > 70%) following the European protocol for subsequent microscopic and metabarcoding analyses (CEN 2014, 2018). Fig. 1 shows the sampling site locations.

DNA extraction and PCR amplification: DNA extraction was done from the pellet obtained following centrifugation of the biofilm (30 min to 17,000 g) using the NucleoSpin Soil kit according to the manufacturer's instructions, as described by Vasselon *et al.* (2017b). A small DNA fragment (312 base pairs in length) of *rbcL* was used for DNA amplification. PCR amplification of the DNA barcode was performed for each sample using a mix of three forward and two reverse primers. The forward primer combined an equimolar mix of Diat_rbcL_708F_1 (AGGTGAAGTAAAA GGTTCTACTTAAA), Diat_rbcL_708F_2 (AGGTGAA GTTAAAGGTTCTACTTAAA) and Diat_rbcL_708F_3 (AGGTGAACTAAAGGTTCTACTTAAA); the reverse primer combined an equimolar mix of R3_1 (CCTTCTAATTTACWACWACTG) and R3_2 (CCTTCTAATTTACWACAACAG) (Vasselon *et al.* 2017b).

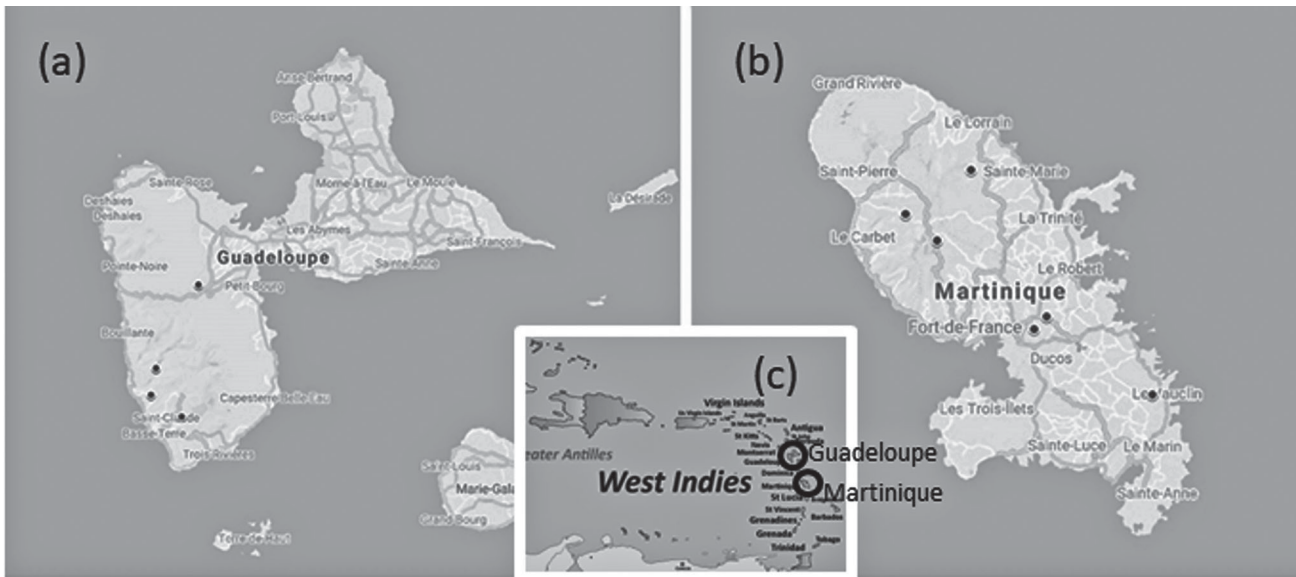


Fig. 1. Maps of Guadeloupe (a), Martinique (b) and general location in the West Indies (c). The location of the sampling sites is marked with black dots, and the locations of the islands are marked with open circles.

Each DNA extract was amplified in triplicate using equimolar mixes of the three forward and two reverse primers. Half the P5 (CTTTCCTACACGACGCTCTTC-CGATCT) and P7 (GGAGTTCAGACGTGTGCTCTTC-CGATCT) Illumina adapters were included to the 5' part of the *rbcL* forward and reverse primers, respectively. Additionally, blank samples using water were run in parallel to check for potential contamination. Amplifications were performed in a final volume of 25 μ L following mix and reaction conditions used previously, the number of amplifications was set to 33 and the conditions of a cycle were as follow: 95°C – 1 min, 54°C – 1 min, 72°C – 1 min (Keck et al. 2018).

High-throughput sequencing and bioinformatics processing: The PCR amplicons were purified and used as templates in a second PCR that used Illumina tailed primers targeting the P5 (CTTTCCTACACGACGCTCTTC-CGATCT) and P7 (GGAGTTCAGACGTGTGCTCTTC-CGATCT) Illumina adapters. Finally, all generated PCR amplicons were indexed and pooled into a single tube. The final pool was sequenced in GeT-Plage (Toulouse, France) Illumina MiSeq platform using the V2 paired-end sequencing kit (250 bp \times 2).

Demultiplexing and adapters' removal were performed by the sequencing platform. Further bioinformatic treatment was performed with R software (3.6.1, R development core team). The software package DADA2 version 1.18.0 (Callahan et al. 2016) was used with parameters adapted to diatom metabarcoding data available on Github (https://github.com/fkeck/DADA2_diatoms_pipeline). The following bioinformatic steps were carried out: (i) primers were removed using cutadapt (version 2.1); (ii) to keep

only good quality sequences (we checked fastqc to have Phred scores above 35), the R1 and R2 reads were truncated to 200 and 170 nucleotides respectively; (iii) only R1 and R2 reads with zero ambiguities and a maximum of two expected errors were kept; (iv) after dereplication, high-quality amplicon sequence variants (ASVs) were selected based on the error rates model and paired reads were merged into one sequence (Tapolczai et al. 2019). The last step (iv) cleaned the data by eliminating chimeric sequences. A total of 1.9 million reads were obtained from a single run of good quality, which comparable to the number of reads obtained in earlier studies (e.g. Rivera et al. 2020), ranging from 2916 to 33192 (average 19533 reads).

To assign taxa to each DNA sequence, the sequences were compared with the reference library Diat.barcode, version 9 (Rimet et al. 2019) using a Naïve Bayesian method with a confidence level of 60% (Youn and Wang 2008). All non-diatom sequences (i.e. not assigned to the 'Bacillariophyta' phylum) were removed. The data were rarified using *rrarefy* function (R package *vegan*, Oksanen et al. 2013) according to the number of reads per sample. The length of the sequences kept after these different selection steps was 263 bp (without primers).

Microscopy: The procedure for slide preparation followed French and European standards (Afnor 2003, 2016). Diatom valves were cleaned using 40% H₂O₂ and 40% HCl. Cleaned valves were mounted in resin (Naphrax[®]). Diatoms were counted as described in the NF T 90-354 standard (Afnor 2016). Specialized floras were used for identification to the lowest taxonomic level, such as diatom floras of the French West Indies (Eulin et al. 2017a, b, c, d, e, f).

Phylogenetic analyses: We constructed a constrained phylogeny by placing the environmental ASVs in a backbone phylogeny. To this end, we first used the alignment available in Diat.barcode v9 (Rimet *et al.* 2019). From this alignment, we selected all long sequences (as noted in Diat.barcode). A total of 3265 sequences were included in the alignment (alignment is available in Diat.barcode file at doi:10.15454/TOMBYZ). The best substitution model was then tested in MEGA7 (Kumar *et al.* 2016). A maximum likelihood (ML) tree was calculated following the best substitution model (GTR + G + I model) with raxmlGUI 2.0 (Stamatakis *et al.* 2005, Silvestro & Michalak 2012). We then added the 100 most abundant ASV sequences a posteriori in the phylogeny. This was also done in raxmlGUI under the ‘enforce constraint menu’ and ‘use multifurcating constraint’. One hundred bootstraps were run, ML and a bootstrap search took 27.26 h with an Intel® Core™ i5-8250U CPU @ 1.60 GHz. The trees presented in the results are parts extracted from this tree and represented using MEGA7.

BLAST analyses: Nucleotide BLAST (Basic Local Alignment Search Tool, blastn) was used to identify homologous sequences (Altschul *et al.* 1990). To this end, we used the local Blast tool proposed in BioEdit version 7.2.5 (Hall 1999). The nucleotide database we used was Diat.barcode v9 (all rbcL sequences), and the maximum e-values used to determine the sequences were e-110.

Correlations between number of ASV and frustule counts: For each unidentified ASV, correlations between abundances of the ASV sequences (reads number) and the abundances of the suspected species counts from the corresponding study sites were carried out using the Pearson correlation coefficient.

Results

According to the degree of difficulty with which we could establish the correspondence between sequence and morphology, three categories were distinguished: (1) simple cases (ASV2-*Ulnaria goulardii*, ASV22-*Nupela* sp. nov., ASV38-*Epithemia* sp. nov.), (2) intermediate cases (ASV11-*Gomphonema bourbonense*), (3) complicated cases (ASV5, ASV8, ASV37-*Nitschia* group). These sequences and their corresponding material (raw material, treated material, slides), as well as their metadata, are registered in the TCC collection and in Genbank (NCBI). All data are also available in Diat.barcode v10.

Simple cases: The first undetermined sequence (ASV2) was identified to family level, namely Fragilariaceae, with a bootstrap value of 97% using DADA2. BLAST identified the sequence as *Fragilaria gracilis* Østrup, with 95% identity. This guided us to the genus we should look for in the LM slides. The position of ASV2 in the phylogenetic tree was inside the *Ulnaria* (Kützing) Compère clade

(Fig. 2). Checking the slides in LM (Figs 3–11) showed that ASV2 corresponded to *Ulnaria goulardii* (Brébisson ex Cleve & Grunow) D. M. Williams, Potapova & C. E. Wetzel, a species recently transferred from *Fragilaria* Lynge to *Ulnaria* (Wetzel *et al.* 2022), and the only *Ulnaria* species observed by LM in the samples. ASV2 read numbers and frustule counts of *U. goulardii* showed a significant correlation ($R^2 = 0.42$, $p < 0.05$). Our results also confirm the transfer to *Ulnaria* proposed by Wetzel *et al.* (2022).

Undetermined sequence ASV22 was also identified to family level, namely Naviculaceae, with a bootstrap value of 99% using DADA2. BLAST identified ASV22 as *Nupela* Vyverman & Compère sp. with 93% identity. Although the bootstrap value (45%) only gave poor support for its position in the phylogeny, this sequence was next to a sequence of *Nupela* (Fig. 12). We therefore correlated the ASV22 read number to the frustule counts of *Nupela* sp., and the results were significant ($R^2 = 0.58$, $p < 0.05$). We therefore concluded that this sequence belongs to a species of *Nupela*. LM observations (Figs 13–53) and SEM observations (Figs 54–59) confirmed the genus affiliation. According to the molecular, LM and SEM observations, this species is new to science and is described below (Taxonomic results).

Undetermined sequence ASV38 was assigned to the genus *Epithemia* Kützing using DADA2. BLAST identified ASV38 with 95% identity to four sequences belonging to *Epithemia gibba* (Ehrenberg) Kützing, *Epithemia hyndmanii* W. Smith and *Epithemia* sp. The position of ASV38 in the phylogeny was inside the *Epithemia* clade (Fig. 60). Earlier identifications carried out for the iconographic atlas of the French West Indies identified a taxon as *Rhopalodia* sp.1 (Eulin *et al.* 2017e). However, this species is now part of *Epithemia* since *Rhopalodia* O. Muller has been merged with *Epithemia* on the basis of morphological and genetic data (Ruck *et al.* 2016). LM (Figs 61–87) and SEM observations (Figs 88–93) of our *Epithemia* sp. present morphological features which differ from closely related known species. ASV38 reads number and frustule counts of *Epithemia* sp. were significantly correlated ($R^2 = 0.1223$, $p < 0.05$). Based on the molecular, LM and SEM results, this species is new to science and is described below (Taxonomic results).

Intermediate case: DADA2 assigned sequence ASV11 to the genus *Gomphonema* Ehrenberg, with 100% bootstrap support. BLAST identified the sequence as *G. bourbonense* E. Reichardt with 97% identity. The position of ASV11 in the phylogenetic tree (Fig. 94) shows that it belongs to the *G. bourbonense* clade, which also includes ASV16. The LM determinations carried out in the earlier study (Eulin *et al.* 2017b) show that two morphologically similar species were identified in the West Indies: *G. bourbonense* and *G. designatum* E. Reichardt. However, we suspected that these identifications were erroneous. We therefore carried out detailed morphometric analyses on

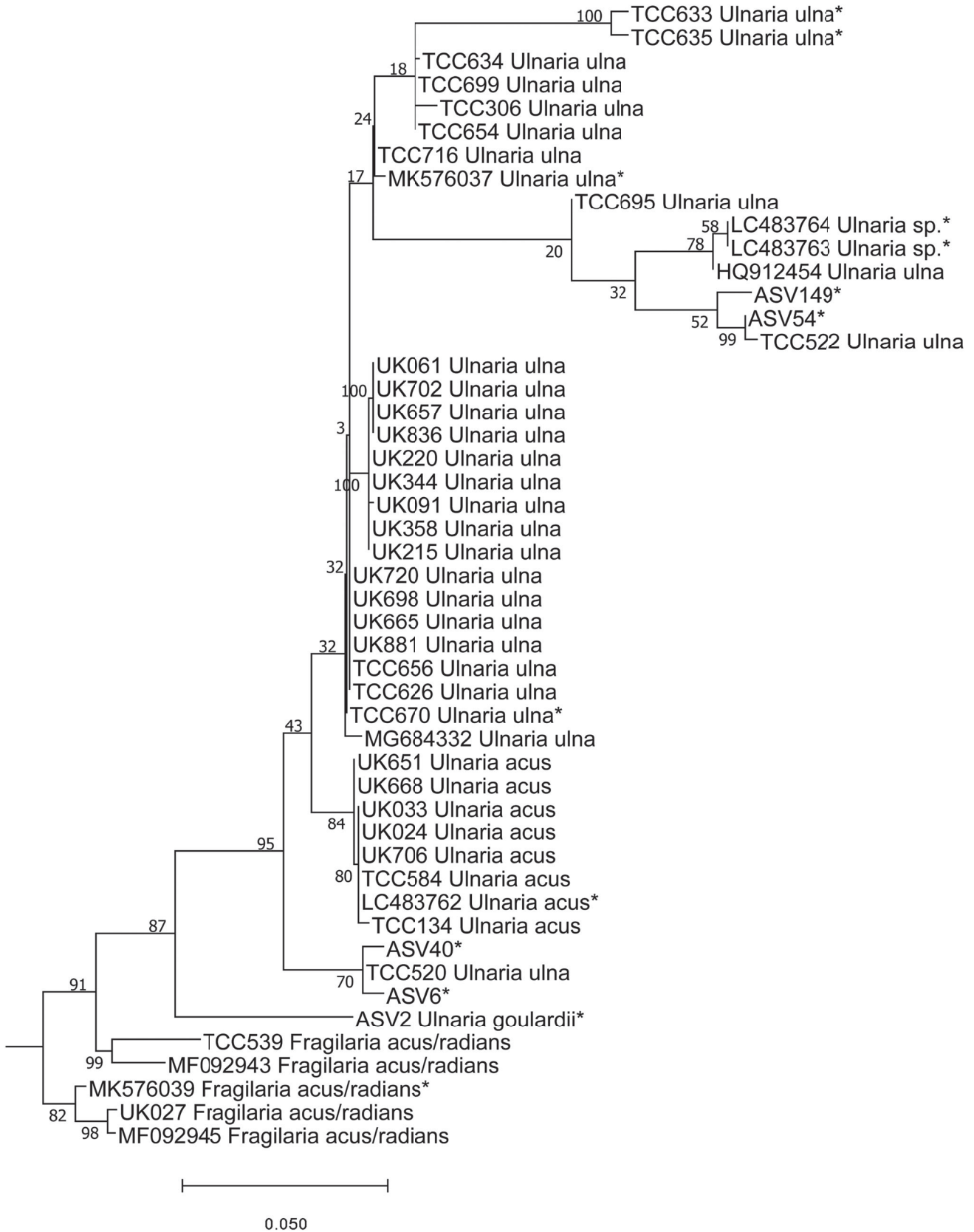
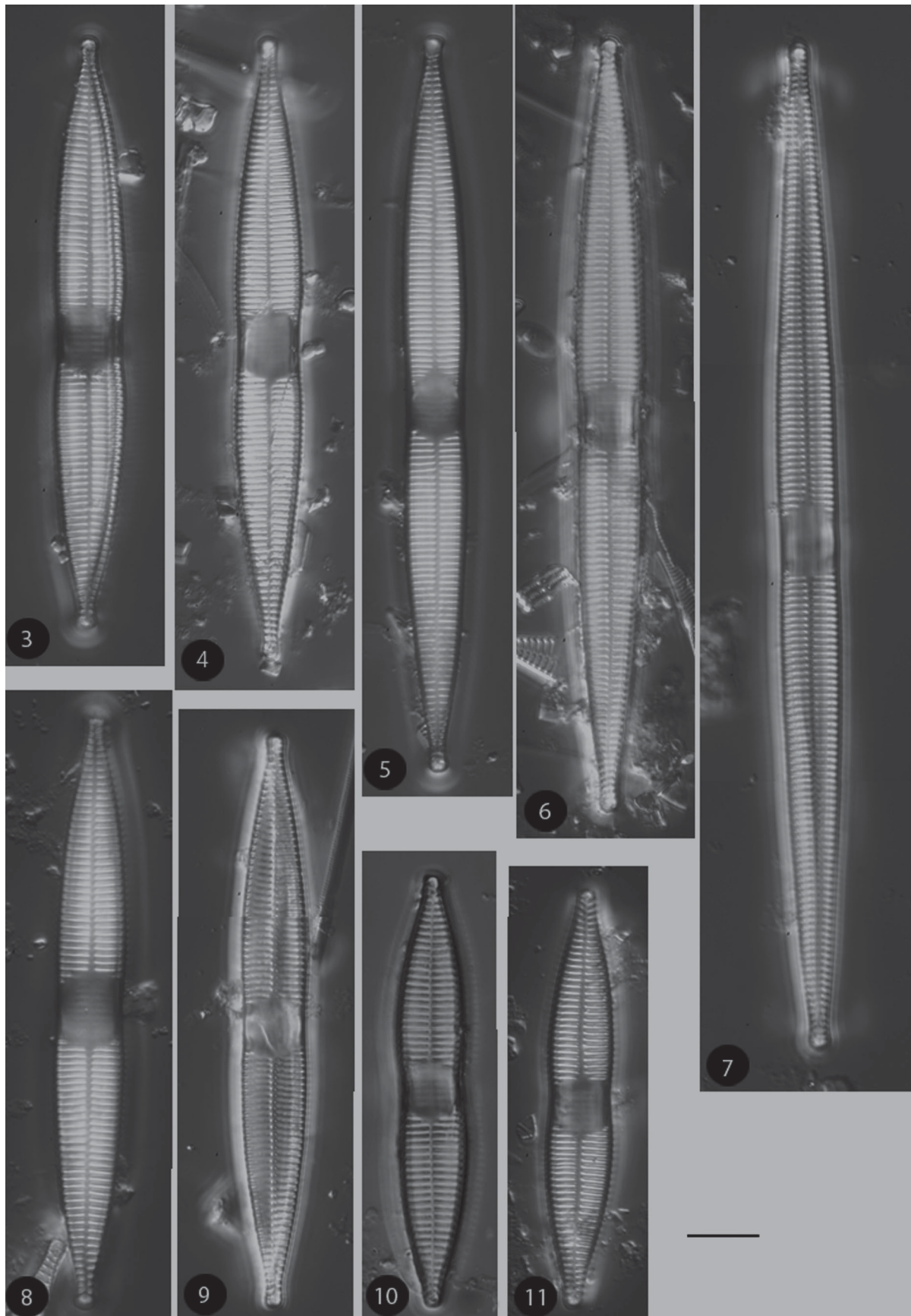


Fig. 2. Phylogenetic position of *Ulnaria gouldarii* (ASV2) in the ML tree. Bootstrap values are given for each node and the scale bar gives the number of substitutions per site. “*” indicates sequences added in the phylogeny using the multifurcating constraint.



Figs 3–11. LM micrographs of *Ulnaria goulardii*. Figs 3–11. Acc. No. TCC1083. From site called Grande Rivière – Vieux habitants amount in Guadeloupe. Scale bar = 10 μ m.

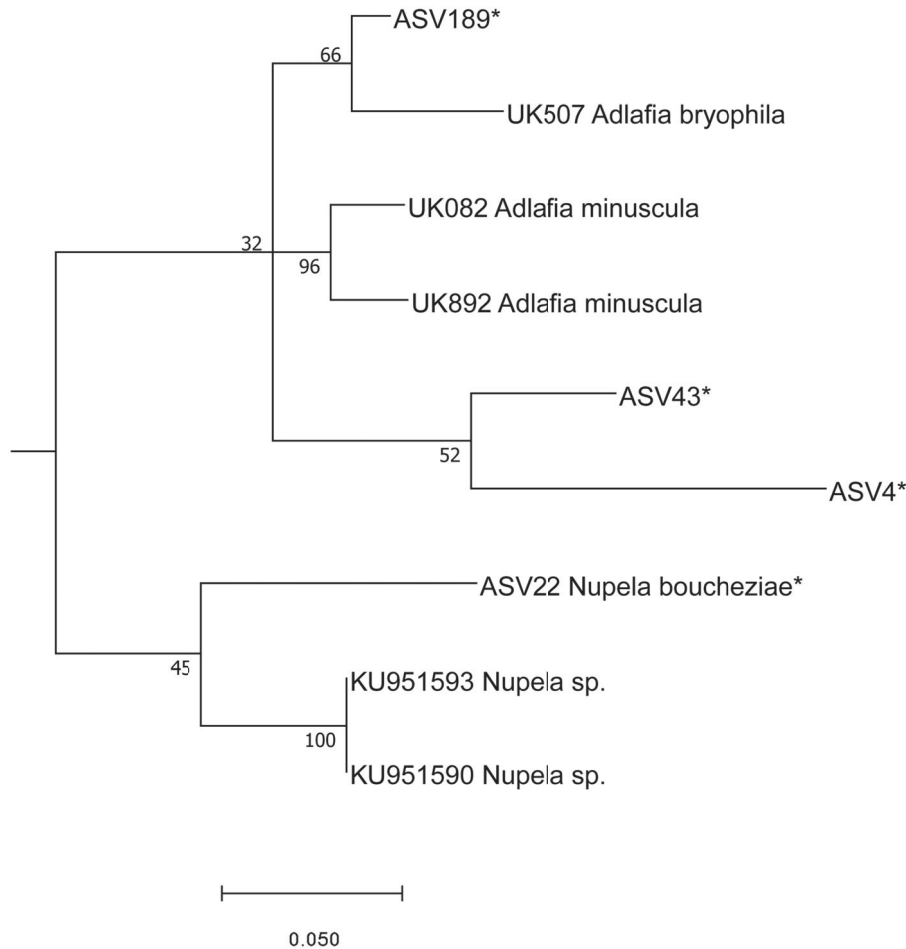
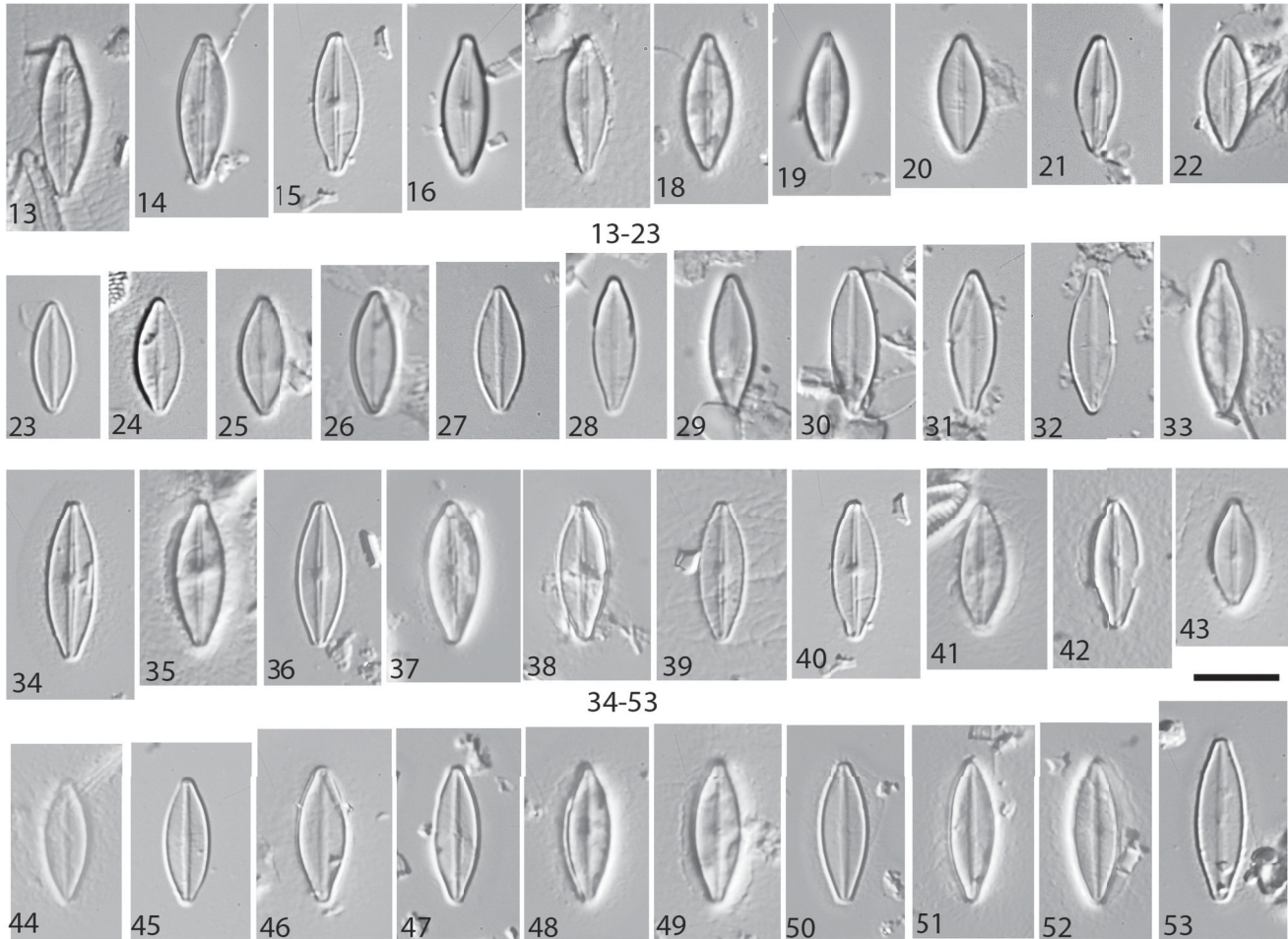


Fig. 12. Phylogenetic position *Nupela boucheziae* sp. nov. (ASV22) in the ML tree. Bootstrap values are given for each node and the scale bar gives the number of substitutions per site. “*” indicates sequences added in the phylogeny using the multifurcating constraint.

the West Indies samples in which these species were formerly identified. Our measurements, given in Table 1, did not fit with *G. designatum* (particularly the ranges for length, width and stria density) but with *G. bourbonense*. Therefore we considered the former determinations of *G. designatum* by Eulin et al. (2017b) should be *G. bourbonense*. Correlation between the numbers of reads of ASV11 plus ASV16 and frustule counts of *G. bourbonense* (to which were added frustule counts previously identified as *G. designatum*) was significant. Therefore, although two species were identified and illustrated in the atlas of these islands (Eulin et al. 2017b), we conclude that they constitute a single species, *G. bourbonense*.

Complex case: A group of sequences were treated together due to their close phylogenetic affiliation. These were ASV5, ASV8, ASV37, ASV57 and ASV150. DADA2 assigned ASV5, ASV8, ASV37, ASV57 and ASV150 to the genus *Nitzschia* Hassall. BLAST gave the following identifications: *Nitzschia amphibia* Grunow (98%) (ASV5), *Nitzschia inconspicua* Grunow (97%) (ASV8),

N. inconspicua (98%) (ASV37), *N. amphibia* (98%) (ASV57) and *N. amphibia* (98%) (ASV150). The position of these five sequences is shown in the phylogenetic tree (Fig. 95); their positions next to *N. inconspicua* and *N. amphibia* are loosely supported. Therefore, all these sequences (ASV5, ASV8, ASV37, ASV57, ASV150) were considered for the correlation between their read numbers and the frustule counts of several *Nitzschia* species. Given their position in the phylogeny, we suspected that they might correspond to *N. inconspicua*, *N. amphibia*, *Nitzschia denticula* Grunow, but also to other morphologically similar taxa illustrated in Eulin et al. (2017e) as *N. frustulum* (Kützing) Grunow; *N. frustulum* forme 2, *N. frustulum* forme 3, *N. sp. 64* and *N. sp. 41*. The correlations (supplementary data 4) show that in several cases, read abundance of an ASV could be correlated to several species abundances identified by LM (e.g. ASV5 correlated with *N. inconspicua* and *N. frustulum*, ASV8 correlated with *N. inconspicua* and *N. amphibia*). Moreover, some ASVs co-occurred (e.g. ASV5 and ASV8), although they were not located in the same phylogenetic



Figs 13–53. LM micrographs of *Nupela boucheziae* sp. nov. Figs 13–33. Type material Acc. No. TCC1086. Figs 13–22. Valves (with raphe) and Figs. 23–33. Valves (without raphe), from site called Blanche-Pont de l’Alma in Martinique. Figs 34–42. Valves (with raphe) and Figs. 43–53 Valves (rapheless), from river Rivière Bras David in Guadeloupe, Acc. No. TCC1088 (paratype). Scale bar = 10 μ m.

clade (ASV5 in *N. amphibia* clade, ASV8 weakly placed in the phylogeny between *N. amphibia*, *N. inconspicua*, *N. denticula*) and were correlated with the same species identified in LM (*N. inconspicua*). For all these reasons, it is impossible to give species names to these ASVs with any certainty.

Taxonomic results

Nupela boucheziae Kochoska, Chardon, Chonova, Keck, Kermarrec, Larras, S.F. Rivera, Tapolczai, Vasselon, Levkov & Rimet sp. nov. (Figs 13–59)

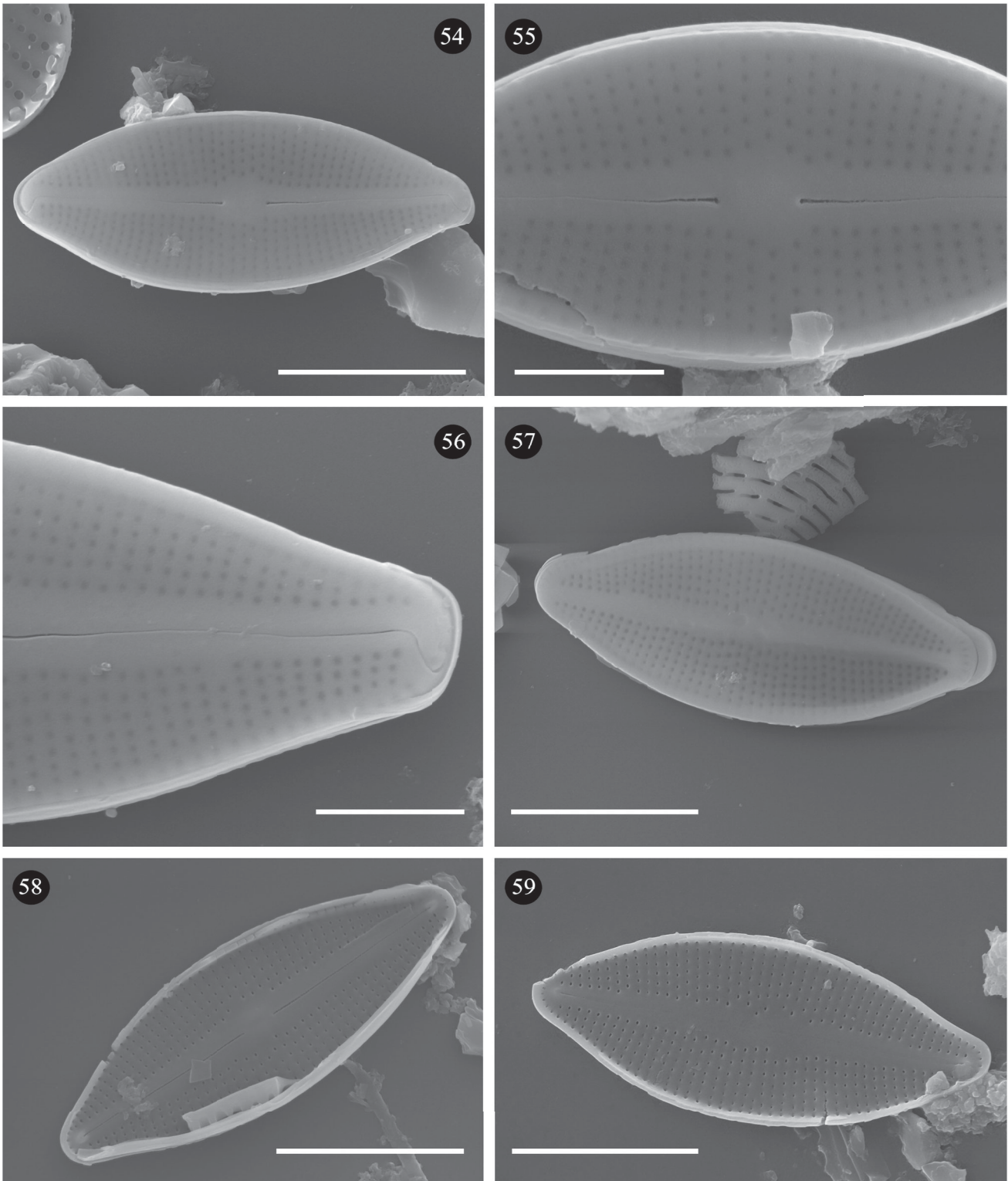
Description: LM (Figs 13–53): Frustules convex, heterovalvar, slightly asymmetric about the apical plane. Valves lanceolate, elliptical-lanceolate with slightly rounded to sub-rostrate apices, 7.0–15.0 μ m long and 4.0–4.5 μ m wide. One valve with long raphe slits, thread-like and almost straight, incomplete on the other valve, reduced to a small helictoglossa, ‘ghost’ full raphe. Axial area linear and narrow in both valvae, central area small to

very small, round to elliptical. Striae and areolae not visible in LM.

Description: SEM (Figs 54–59): Heterovalvar frustules. Proximal raphe ends externally expanded and internally simple (Figs 55, 58). Terminal raphe ends curved externally to the same side of the valve and internally ending in small helictoglossae (Figs 54, 56 and 58, 59). Reduced raphe valve with a smooth axial area without depressions (Figs 57, 59). Transapical striae are slightly radial to parallel towards the apices, 30–40 in 10 μ m, composed of continuous lines of areolae, ca. 50 in 10 μ m. Outer openings of areolae are round and occluded by a delicate hymen. Inner openings of areolae larger than the outer, round to oval.

Type: France, Blanche-Pont de l’Alma in Martinique, biofilm, collection date: 12.05.2018; Leg. Anne Eulin, Estelle Lefrançois; Coordinates: – 61.08895606 latitude, 14.70644106 longitude.

Holotype slide and treated material: Accession No. PC0643142 (Museum National d’Histoire Naturelle, Paris, France).



Figs 54–59. SEM micrographs of *Nupela boucheziae* sp. nov. from type material TCC1086. Figs 54–57. External valve view. Figs. 58–59 Internal valve view. Scale bars = 5 μ m (54, 57–59). Scale bars = 2 μ m (55, 56).

Isotype slide and treated material: Accession No. MKNDC 14432 (Institute of Biology, Skopje, Republic

of North Macedonia). Slide and treated material TCC1086 (Thonon Culture Collection).

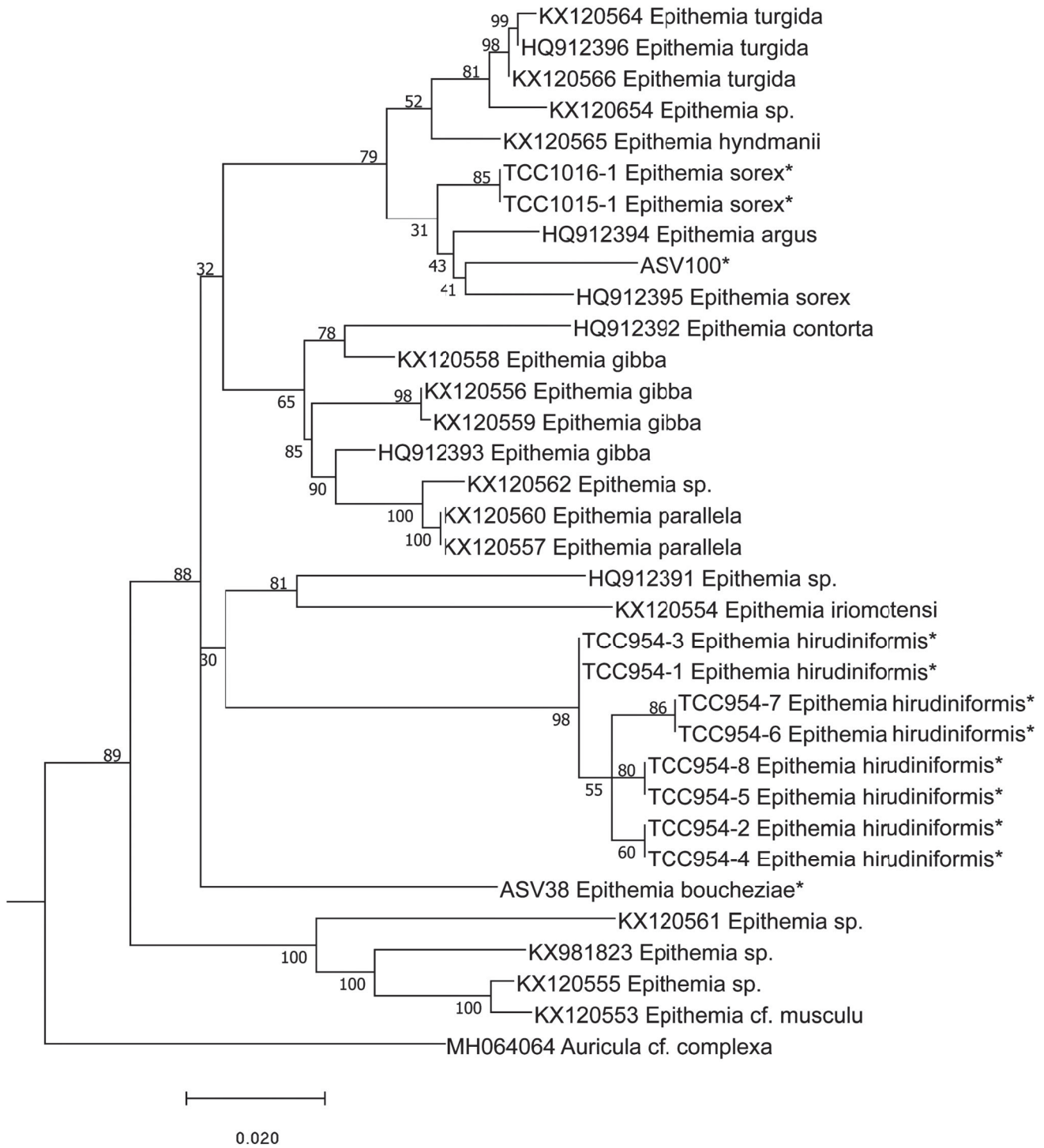
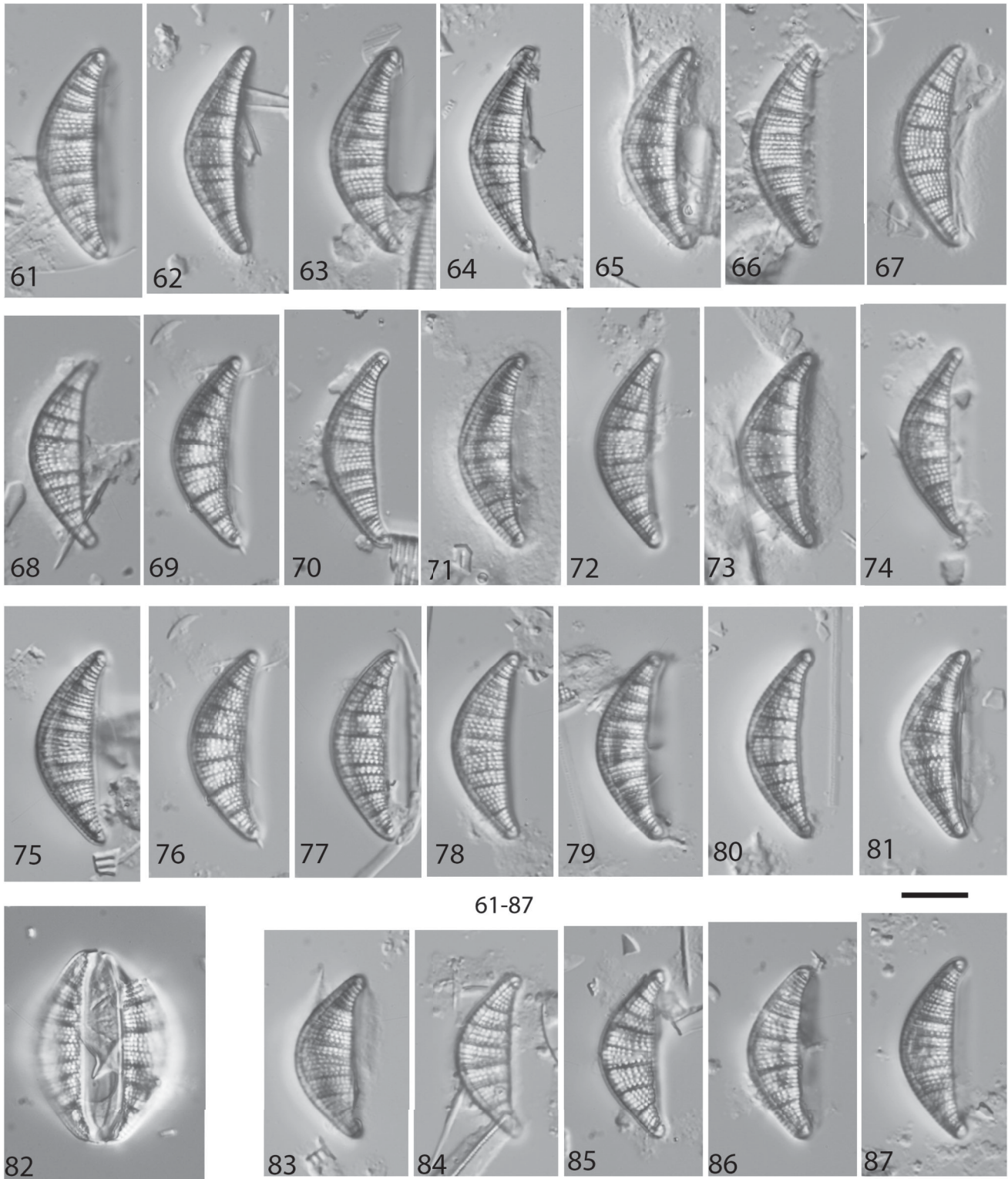


Fig. 60. Phylogenetic position *Epithemia boucheziae* sp. nov. (ASV38) in the ML tree. Bootstrap values are given for each node and the scale bar gives the number of substitutions per site. "*" indicates sequences added in the phylogeny using the multifurcating constraint.

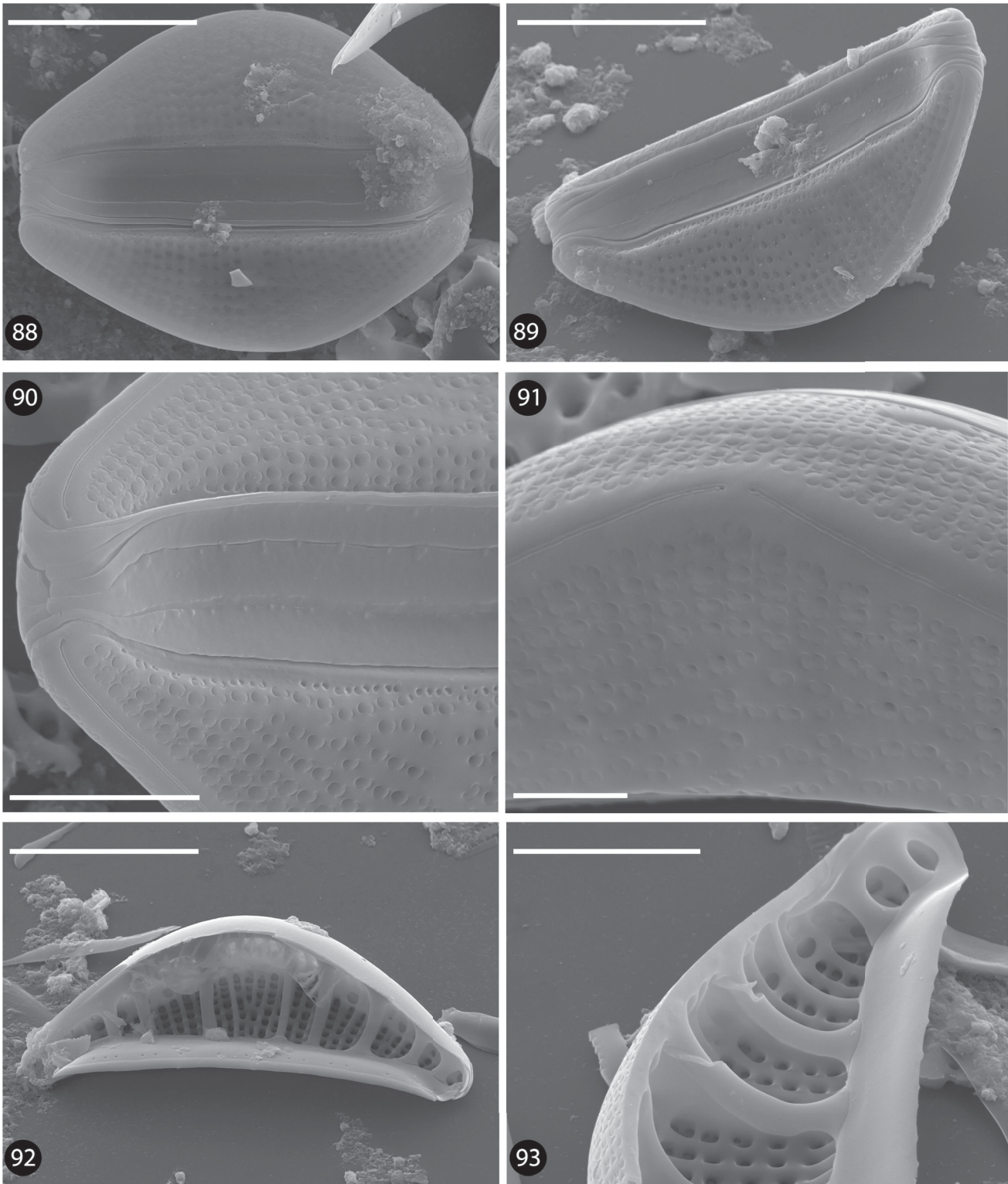
Etymology: This species is dedicated to Dr. Agnès Bouchez, who made major contributions to the development of metabarcoding and diatom science.

Taxonomic remarks: Based on the valve shape, *N. boucheziae* is comparable to several *Nupela* species (see Table 2). *Nupela boucheziae* is similar to *N. praecipuoides*

(Tremarin & T. Ludwig (Tremarin *et al.* 2015) but *N. praecipuoides* has one valve with long raphe slits while the other valve is araphid. Clear differences between *N. boucheziae* and *N. praecipuoides* can also be seen in that the axial area is linear and narrow on the raphid valve, but lanceolate, smooth or generally with irregular



Figs 61–87. LM micrographs of *Epithemia boucheziae* sp. nov. Figs 61–87. Type material Acc. No. TCC1089. Figs 61–87. from river Rivière Bras David in Guadeloupe. Scale bar = 10 µm.



Figs 88–93. SEM micrographs of *Epithemia boucheziae* sp. nov. from type material TCC1089. Figs 88–91. External valve view. Figs 92–93. Internal valve view. Scale bars = 10 μ m (88, 89, 92). Scale bars = 5 μ m (90, 93). Scale bars = 2 μ m (91).

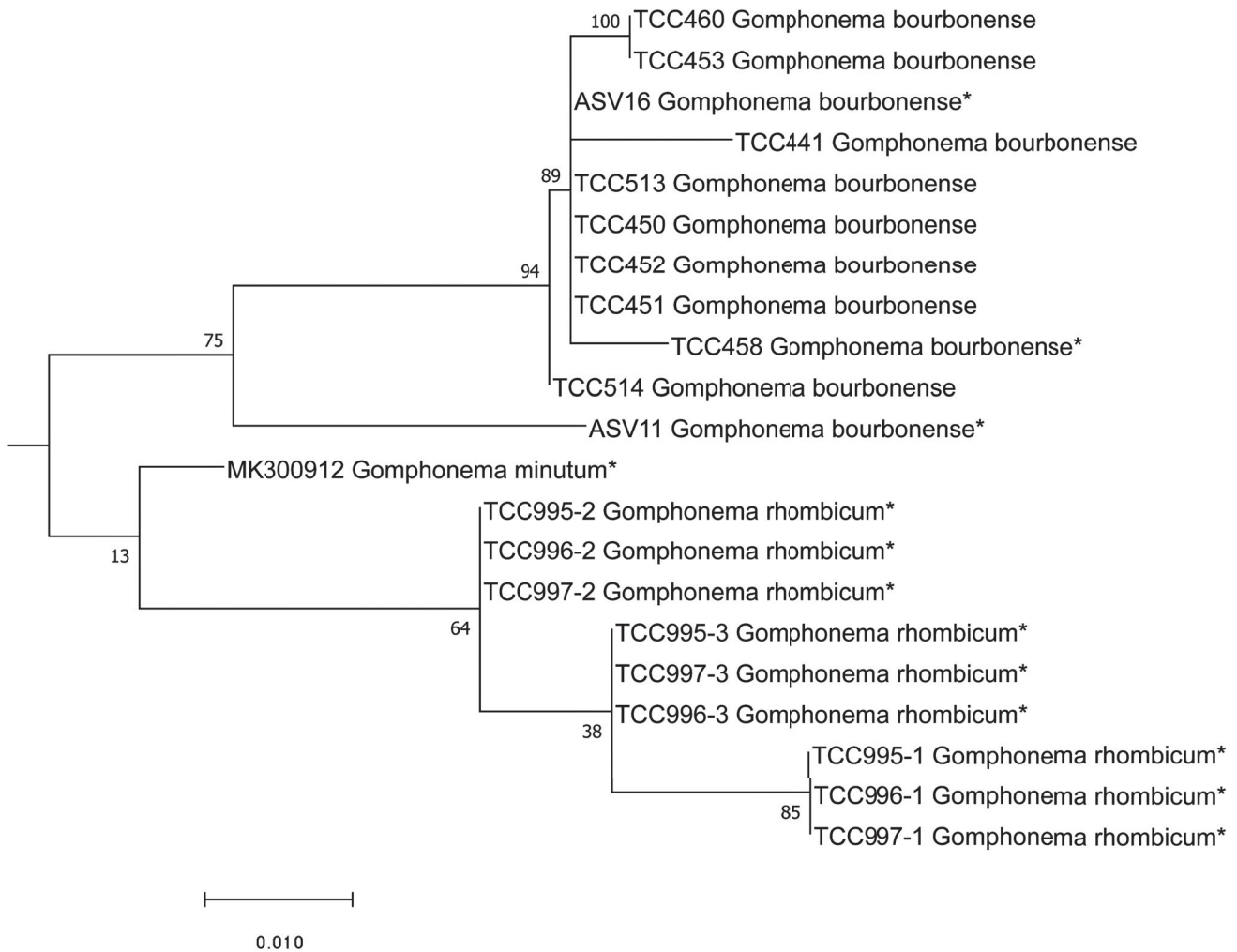


Fig. 94. Phylogenetic position *Gomphonema bourbonense* (ASV11 and ASV16) in the ML tree. Bootstrap values are given for each node and the scale bar gives the number of substitutions per site. "*" indicates sequences added in the phylogeny using the multifurcating constraint.

depressions on the araphid valve that may or not be visible in LM. *Nupela praecipuoides* also has much lower areola and stria densities. Specimens similar to *N. praecipuoides* were recorded by Rumrich et al. (2000) as *N. spec. cf. praecipua* in Ecuador, and in south Brazil as *N. praecipua* (E. Reichardt) E. Reichardt by Schneck et al. (2008), Tremarin et al. (2009) and Moresco et al. (2011). *Nupela praecipuoides* was also recorded in rivers of the Atlantic forest in southern Brazil (Tremarin et al. 2015).

Nupela boucheziae is similar to *N. praecipua* described from Mexico (Reichardt 1988). Both species share a similar valve outline, but the striae and areolae of *N. praecipua* are coarser (32–36 striae in 10 μm , 30–35 areolae in 10 μm) than those of *N. boucheziae*: this is easily seen under LM. Furthermore, *N. praecipua* has smaller valves (length: 8.0–13.5 μm) and deep depressions in the axial area of the araphid valve, which is not the case for *N. boucheziae*. *Nupela praecipua* has slightly convergent striae at the apices, unlike *N. boucheziae* (Reichardt

1988, Rumrich et al. 2000). Some similarity was observed between *N. boucheziae* and *N. chilensis* (Krasske) Lange–Bertalot (Lange–Bertalot et al. 1996) in valve outline and striation pattern. However, *N. chilensis* has larger valves (length: 16–26, width: 5–7 μm), a wider central area, long raphe slits on both valves, and lower stria density (30–32 in 10 μm) than *N. boucheziae*. *Nupela difficilis* Straube, Tremarin & T. Ludwig is mainly characterized by its lanceolate valve outline, subrostrate apices, and asymmetric central area, as well as the straight, interior, proximal raphe ends (Tremarin et al. 2015). *Nupela difficilis* has a similar valve outline and size to *N. boucheziae*. However there are clear differences with a raphe on both valves and strongly convex valve margins with its asymmetric central area, reaching the valve margin on one side.

Nupela decipiens (Reimer) Potapova is comparable in size and stria density to *N. boucheziae* (Potapova 2013). *Nupela decipiens* can be differentiated by the narrowly rostrate to subrostrate apices, the size and shape of the central

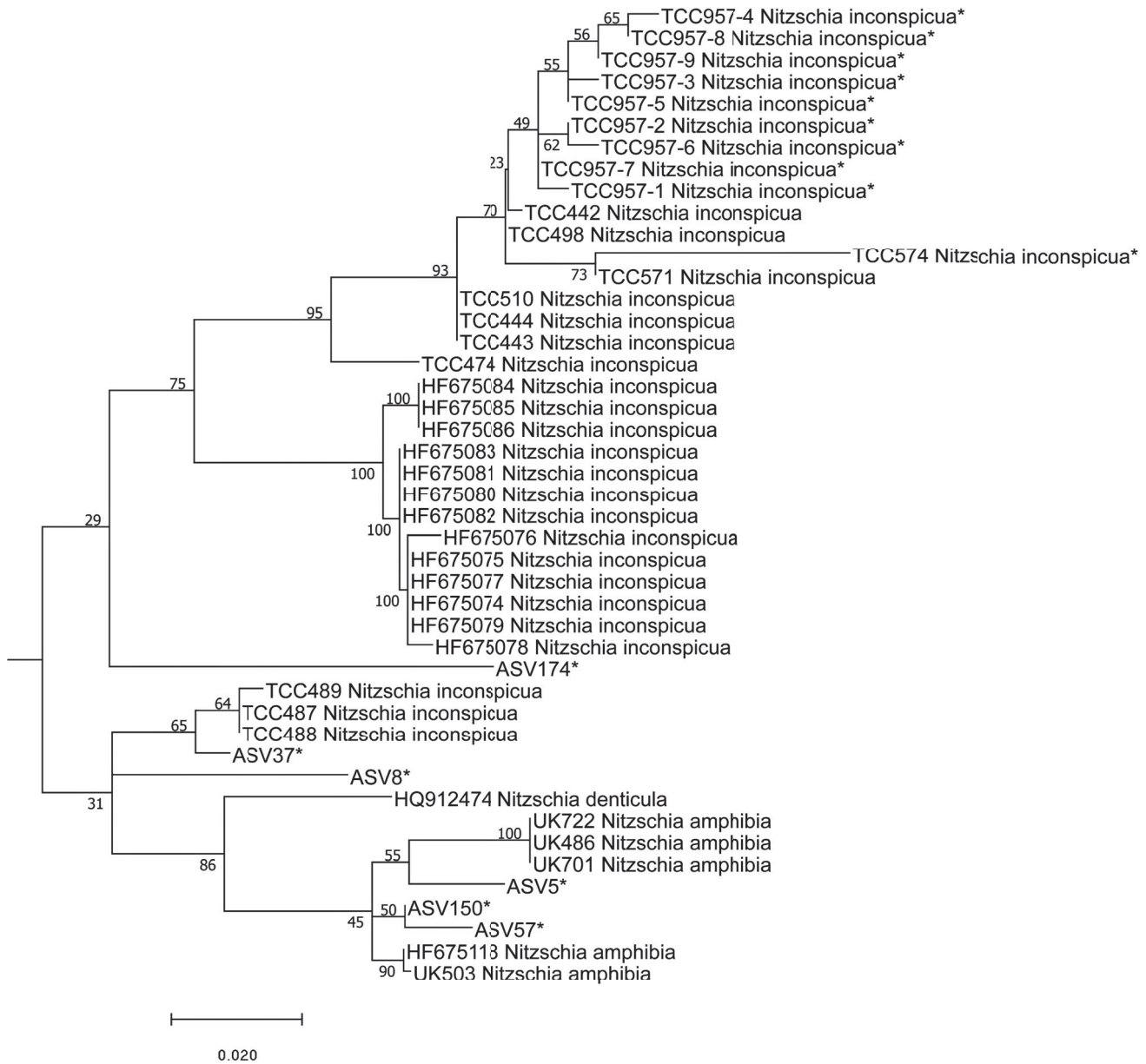


Fig. 95. Phylogenetic position of undetermined ASV (ASV5, ASV8, ASV37, ASV57, ASV150) of the *Nitzschia* complex in the ML tree. Bootstrap values are given for each node and the scale bar gives the number of substitutions per site. "*" indicates sequences added in the phylogeny using the multifurcating constraint.

area (in raphe widely rounded, in araphid valve widely lanceolate) not reaching the valve margin. The axial area also differs between the two species: lanceolate to widely lanceolate, with irregular external depressions, the longitudinal depression in the araphid valve resembling a raphe under LM. *Nupela neglecta* Ponader, Lowe & Potapova, and *N. boucheziae* share similar valve outlines (Potapova *et al.* 2003); however, *N. neglecta* has slightly protracted apices and quite different valve margins: one valve is convex to parallel in the middle, the other slightly concave and often slightly asymmetrical about apical and transapical planes. *Nupela neglecta* has raphe slits on both valves (one raphe shorter than the other),

and a higher stria density (40–48 in 10 μ m) compared to *N. boucheziae*. *Nupela jahniae-reginae* Lange-Bertalot & Metzeltin (Rumrich *et al.* 2000) also has a comparable valve shape to *N. boucheziae*. (elliptical lanceolate to lanceolate). But there are differences between the two species: *N. jahniae-reginae* has obtuse, slightly protracted apices, convex to slightly parallel in the middle valve margins, with a longer raphe on the short raphe valve and lower valve width (3.0–4.0 μ m in *N. jahniae-reginae*) and higher stria density (ca. 50 in 10 μ m in *N. jahniae-reginae*).

Ecological remarks: This species is abundant in the West Indies (Martinique and Guadeloupe). The type locality is

Table 1. Morphological comparison between *Gomphonema bourbonense* and *Gomphonema designatum*.

Morphological characteristics	<i>Gomphonema bourbonense</i> E. Reichardt	<i>Gomphonema designatum</i> E. Reichardt	<i>Gomphonema bourbonense</i> (this study)
Valve outline	Linear-elliptic to linear-lanceolate	Moderately heteropolar	Linear to linear-lanceolate
Apices	Rounded, never rostrate	Rounded to pointed	Rounded
Length	9.4–28 µm	22–45 µm	13.0–25.0 µm
Width	3.3–4.7 µm	4–5.5 µm	3.40–4.5 µm
Stria density	10.5–13/10 µm	10–12/10 µm	12–13/10 µm
Stria orientation	Parallel to slightly radiate	Strong, spaced parallel becoming radiate in larger specimen	Parallel to slightly radiate
Central area	Rectangular with 1 stigma	Poorly individualized, bordered by 2 shorter striae	Rectangular with 1 stigma

situated in the upstream stretch of the Alma river. It is characteristic of low nutrients and organic matter (Eulin et al. 2017d).

Slides are deposited at the National Museum of Natural History (MNHN) in Paris, France and the Macedonian National Diatom Collection (MKNDC) at the Institute of Biology, Faculty of Natural Sciences, Skopje, Republic of North Macedonia.

Epithemia boucheziae Kochoska, Chardon, Chonova, Keck, Kermarrec, Larras, S.F. Rivera, Tapolczai, Vasselon, Levkov & Rimet sp. nov. (Figs 61–93)

Description. LM (Figs 61–87): Frustules lanceolate to linear-elliptic in girdle view. Valves semi-elliptical to almost triangular, dorsal edge of valve convex, ventral edge slightly concave. The valve ends slightly protracted and curved towards the ventral edge of the valve. Valves 17.0–23.0 µm long and 6.5–8.0 µm in width. The raphe is located in a channel along the dorsal edge of the valve. Areolae visible with LM, ca. 16 in 10 µm.

Description. SEM (Figs 88–93): Externally, proximal raphe ends almost straight, slightly expanded to droplet shaped. Terminal raphe ends curved ventrally. Internal raphe endings are simple. Raphe opens internally into a canal with small round holes (portulae) lying between the major transapical ribs (Fig. 93). Striae coarsely punctate and strongly radiate. Externally, striae are composed of complex areolae: near the ventral side these are composed of two opposed ‘C’ shaped slits, and near the dorsal side of four opposed ‘C’ shaped slits, giving a flower-like aspect (Fig. 91). Discontinuous stria pattern with several missing areolae in the middle part of the valve (Fig. 89). Striae uniseriate, 16–18 in 10 µm. Primary fibular costae 3–4 in 10 µm with usually 4–6 striae between two costae (Figs 92, 93).

Type. France: Rivière Bras David in Guadeloupe, biofilm, collection date: 16.04.2019; Leg. Anne Eulin,

Estelle Lefrançois; Coordinates: – 61.67076118 latitude, 16.19470576 longitude.

Holotype slide and treated material: Accession No. PC0643143 (Museum National d’Histoire Naturelle, Paris, France).

Isotype slide and treated material: Accession No. MKNDC 14433 (Institute of Biology, Skopje, Republic of North Macedonia). Slide and treated material TCC1088 (Thonon Culture Collection).

Etymology: This species is dedicated to Dr. Agnès Bouchez, on behalf of her close colleagues and former students who all enjoyed working with her.

Taxonomic remarks: Based on morphological features, *E. boucheziae* is similar to several species previously placed in *Rhopalodia* (Table 3). Valve shape of *E. boucheziae* is similar to *R. michelorum* Krammer, which has a nearly straight ventral margin, and quite distinct ends, which are narrowly protracted, frequently capitate, and bent ventrally (Krammer 1988). However, *R. michelorum* has bigger valves (length: 17–50 µm, width: 6–10 µm), its proximal raphe ends are curved to the same side, hook-like (*E. boucheziae* has straight proximal raphe ends), and the striae are composed of single rows of areolae. The stria pattern is discontinuous with several missing areolae in the middle part of the valve. Stria density is higher in *R. michelorum* (19–24 in 10 µm) than *E. boucheziae* (16–18 in 10 µm) and the primary fibular costae of *R. michelorum* are very stout and more distantly spaced (1.5–2.8 in 10 µm, Table 3).

Epithemia boucheziae is usually smaller than *Rhopalodia gibberula* (Ehrenberg) O. Müller, however, the smaller individuals of *R. gibberula* are similar to *E. boucheziae*. The difference can be seen in the valve outline, which is linear-elliptic to broad-elliptical in *R. gibberula* and hardly or not retracted in the middle, and the sides are massiform to strongly convex. Frustules of *R. gibberula* are

Table 2. Morphological features and measurements of *Nupela boucheziae* sp. nov. and similar species. n.a.: no data available.

Species	Valve outline	Apices	Valve margins	Central area	Raphe	Axial area	Stria orientation	Length (µm)	Width (µm)	Areola density (in 10 µm)	Stria density (in 10 µm)	Ref.
<i>Nupela boucheziae</i> sp. nov.	Elliptical-lanceolate	Rounded to sub-rostrate	Convex	Small to very small, round to elliptical	Thread like and almost straight, incomplete on one valve, reduced to small helictoglossa, 'ghost' full raphe	Narrow on both valves	Radial to parallel towards the ends	7.0–15	4–4.5	ca. 50	30–40	1
<i>Nupela praecipuoides</i>	Lanceolate	Slightly protracted to subrostrate	Convex	Rounded	One valve with long raphe slits, other valve araphid	Raphid valve - linear and narrow. Araphid valve - lanceolate, smooth or with irregular depressions	Radial to parallel at the apices	9.0–21.5	4.0–5.5	35–39	36–38	2
<i>Nupela praecipua</i>	Broadly lanceolate	Obtuse ends	Convex	Unmarked	Filiform. Slightly curved. One valve with complete other with rudimentary raphe	Wide, with irregular boundaries	Radial throughout valve, slightly convergent at the apices	8–13.5	4–5	35–43	30–39	3, 4
<i>Nupela chilensis</i>	Linear-elliptic	Cuneiform narrowed, bluntly rounded	Convex	Quite big and roundish, well defined	Filiform on both valves	Narrower to very broad, lanceolate, widening towards the middle		16–26	5–7	n.a.	32	5, 6
<i>Nupela difficilis</i>	Lanceolate	Subrostrate	Strongly convex	Asymmetric, unilaterally reaching valve margin	One valve with long raphe slits. Other valve with shorter raphe slits	Lanceolate	Transapical, slightly radiate	8.8–14.8	4.1–5.9	48–50	40	2
<i>Nupela decipiens</i>	Lanceolate	Narrowly rostrate to subrostrate	Convex to parallel in middle	Raphid valve - broadly rounded. Araphid - broadly lanceolate, not reaching valve margin	One valve with long raphe slits and other valve araphid	Raphid valve-lanceolate. Araphid valve - broadly lanceolate, externally ornamented with irregular depressions	Transapical radiate.	8.9–18.5	4.1–5.9	52	36–40	6, 7
<i>Nupela neglecta</i>	Lanceolate to elliptical-lanceolate	Slightly protracted	One valvae convex to parallel in middle. Other slightly concave or slightly asymmetrical in apical and transapical planes	Small, round or elliptical	One valve - long raphe slits. Other valve - distinctly shorter raphe slits. Widely separated central endings	Linear-lanceolate	Radiate, becoming parallel or slightly convergent near the poles	3–15	2.6–4.5	n.a.	40–48	8, 9
<i>Nupela jahninae-reginae</i>	Elliptical-lanceolate to lanceolate	Obtuse, slightly protracted	Convex to slightly parallel in middle	Wide, linear- elliptic	Filiform on both valves, one shorter than other	Linear, narrow	Radiate to convergent at apices	9–14	3–4	n.a.	ca. 50	10

1: This study

2: Tremarin et al. 2015.

3: Reichardt 1988.

4: Rumrich et al. 2000.

5: Krasske 1939.

6: Lange-Bertalot & Moser 1994.

7: Potapova 2013.

8: Reimer 1966.

9: Potapova et al. 2003.

10: Rumrich et al. 2000.

Table 3. Morphological features and measurements of *Epithemia boucheziae* sp. nov. and similar species. n.a.: no data available.

Species	Valve outline	Apices	Valve margins	Raphe	Striae	Length (µm)	Width (µm)	Costa density (in 10 µm)	Stria density (in 10 µm)	Stria number between two costae	Ref
<i>Epithemia boucheziae</i> sp. nov.	Semi-elliptical	Rounded, slightly curved	Dorsal edge, clearly convex, ventral edge slightly concave	Proximal raphe ends almost straight, slightly expanded-drop shaped. Terminal raphe ends curved to same side of valve	Uniseriate	18–23.0	5.5–7.0	3–4	16–18	4–6	1
<i>Rhopalodia michelorum</i> Krammer	Strongly dorso-ventral	Quite distinct, narrow, protracted, frequently capitate and bent ventrally	Dorsal convex and ventral nearly straight	Proximal raphe ends curved to the same side	Single rows of areola	17–50	6–10	1.5-2.8	19–24	> 8	2
<i>Rhopalodia gibberula</i> (Ehrenberg) O. Müller	Linear-elliptical to broad-elliptical	Broadly rounded to truncate	Dorsal side strongly convex, ventral side strongly concave in large forms, in smaller cells less curved	Proximal raphe ends with fissure in central portion	The areolae are provided with tube-like processes and bordered by C-shaped foramina	25–100	5–12	3–10	12–19	2–8	3, 4
<i>Rhopalodia gibberula</i> (Ehrenberg) O. Müller	Lanceolate to linear-elliptical, retracted in middle	Narrow, protracted, usually capitate and bent ventrally	Dorsal side convex, ventral side less curved, parallel to slightly concave	Proximal raphe ends with fissure in central portion	n.a.	30–33.5	7–8	6–7	20–28	n.a.	5
<i>Rhopalodia gibberula</i> var. <i>miniuens</i> O. Müller	Elliptical, semicircular	Protracted and bent ventrally	Dorsal side very strongly convex, ventral side straight	n.a.	n.a.	26.5	10	5–6	18–20	n.a.	5

(Continued).

Table 3. Continued

Species	Valve outline	Apices	Valve margins	Raphe	Striae	Length (μm)	Width (μm)	Costa density (in 10 μm)	Stria density (in 10 μm)	Stria number between two costae	Ref
<i>Rhopalodia gibberula</i> var. <i>succincta</i> (Brebisson) Fricke	Linear-elliptical	Broadly rounded slightly curved	Dorsal convex and ventral nearly straight	n.a.	n.a.	21–22	5–6	4–5	17–21	n.a.	5
<i>Rhopalodia gibberula</i> var. <i>Van Heurckii</i> O. Müller	Linear-elliptical, retracted in the middle	Narrow, prolonged and bent ventrally	Dorsal convex and ventral concave to straight	n.a.	n.a.	40–45	9.5	4	18–20	n.a.	5
<i>Rhopalodia musculus</i> (Kutzing) O. Müller	Semi-elliptic	Narrowly rounded	Dorsal strongly convex and ventral side concave	Proximally, raphe slits bend ventrally ending in round central pores	Uniseriate	12–80	10–16	3–5	15–20	2–4	6
<i>Rhopalodia acuminata</i> Krammer	Sickle-shaped	Acutely rounded and narrow	Strongly convex dorsally and weakly concave ventrally	n.a.	Uniseriate becoming multiseriate	22–112	7.5–11	4–6	16–19	n.a.	6, 7
<i>Rhopalodia brebissonii</i> Krammer	Broad-elliptical	Protracted and ventrally bent	Strongly convex dorsal margin, straight or slightly concave ventral margin	Almost straight	Double rows of puncta on both sides of raphe canal; single on rest of valve	15–40	5–8.5	3.5–6	17–22	2–5	6

1: This study.

2: Krammer K. 1988.

3: Ehrenberg C.G. 1843.

4: Lange-Bertalot H., Krammer K. 1987.

5: Bourrelly P. & Manguin E. 1952.

6: Krammer K. & H. Lange-Bertalot 1988.

7: Lange-Bertalot H. & Krammer K. 1987.

sickle shaped, the dorsal side strongly convex, the ventral side strongly concave in large forms, almost parallel to the dorsal side, in smaller ones the ventral side is less curved. The difference between both species is easily noted in the proximal raphe ends: *R. gibberula* has a fissure in the central portion and areolae with tube-like processes and bordered by C-shaped foramina (Lange–Bertalot & Krammer 1987). A population of *R. gibberula* was recorded from freshwater assemblages in Guadeloupe by Bourrelly & Manguin (1952) (Table 2, ref. 5), but this has bigger valves, higher stria density, and a different valve outline. Bourrelly & Manguin (1952) observed different varieties of the ‘gibberula’ group in freshwaters of Guadeloupe: *R. gibberula* var. *miniuens* O. Müller, *R. gibberula* var. *succincta* (Brébisson) Fricke, *R. gibberula* var. *vanheurckii* O. Müller and several others. *Rhopalodia gibberula* var. *miniuens* can be differentiated by its elliptical, semicircular valve outline, the apices are protracted and bent ventrally and the dorsal side is strongly convex, but the ventral side is straight. This species is also wider, with higher costa and stria densities. *Rhopalodia gibberula* var. *succincta* has similarities with *E. boucheziae* in valve shape and apices, but it is smaller and has higher costa and stria densities. *Epithemia boucheziae* is comparable to *R. gibberula* var. *vanheurckii* but differences are clear in the valve outline (linear-elliptical, retracted in the middle in *R. gibberula* var. *vanheurckii*), and *R. gibberula* var. *vanheurckii* is narrower, has more prolonged apices, a larger frustule and a higher stria density (Table 3).

Epithemia musculus Kützing is similar to *E. boucheziae*, characterized by broadly elliptical frustules in girdle view, usually with rounded apices (Krammer 1988). Valves have strongly convex dorsal margins and straight ventral margins, apices are bent ventrally and rounded. This valve outline is different from that of *E. boucheziae*. *Epithemia musculus* is also larger, with more densely arranged, uniseriate striae comprising contrastingly structured areolae. The latter are stout, large and very distinct, less than 15/10 µm, with multiple lips in the foramina, which are unique to *E. musculus* (Table 3).

Rhopalodia acuminata Krammer is morphologically similar to *E. boucheziae*. Its most specific differences are in its shape: the frustule is sickle-shaped, with a strongly convex dorsal margin and weakly concave ventral margin. The areolae in *R. acuminata* are uniseriately arranged with some double rows beside the raphe canal. Areolae are occluded externally by circular or C-shaped slits in *E. boucheziae*. *Rhopalodia acuminata* is also larger than *E. boucheziae*.

Rhopalodia brebissonii Krammer also resembles *E. boucheziae*. This species is characterized by broadly elliptical frustules and valves with a strongly convex dorsal margin, straight or slightly concave ventral margin, and protracted and ventrally bent apices (Krammer 1988). The main differences between *R. brebissonii* and *E. boucheziae*

are in their stria structure, with striae in *R. brebissonii* composed of double rows of areolae on both sides of the raphe canal and a single row on the rest of the valve, while *E. boucheziae* has striae composed of complex areolae. In addition, *R. brebissonii* has larger valves and higher stria density (17–22 in 10 µm) compared to *E. boucheziae*.

Slides are deposited at the National Museum of Natural History in Paris (MNHN) France and the Macedonian National Diatom Collection (MKNDC) at the Institute of Biology, Faculty of Natural Sciences, Skopje, Republic of North Macedonia.

Discussion

Rimet et al. (2018) proposed a methodology to use HTS sequencing data of environmental samples to define barcodes for species which have no sequence in reference barcoding libraries. When we applied this approach to our samples from the West Indies (Guadeloupe, Martinique), the applicability of this approach was variable. We distinguished three categories depending on the level of support between the criteria used in this approach, from simple to complex.

Simple cases: The first group is a category for which correspondence between an unidentified sequence and a morphological specimen can easily be established. These cases were simple because no closely related species were present in the samples. In some cases, the morphological form could be easily assigned to an existing species (*U. gouldarii*), in the other cases it concerned species new to science (*N. boucheziae*, *E. boucheziae*).

The first case (ASV2) concerned a sequence that had the highest number of ASV reads in the rivers of the West Indies. However, it was only identified to the family level (*Fragilariaceae*) with Diat.barcode version 9 because the latter did not contain any similar reference barcode to allow correct naming. Based on molecular and LM data, we could identify this sequence as *U. gouldarii*, a species recently transferred from *Fragilaria* to *Ulnaria* (Wetzel et al. 2022). No similar or related species were observed with LM or in the sequencing data, which made this case straightforward.

Another case was a sequence (ASV22) assigned to the genus *Nupela* on the basis of molecular data. As LM and SEM observations did not match any of the most morphologically similar described species in this genus (Potapova et al. 2003, Potapova 2011, Tremarin et al. 2015), this taxon was described as a new species, *N. boucheziae*.

The last simple case concerned another new species belonging to *Epithemia*. This taxon had previously been observed in the West Indies islands and referred to as *Rhopalodia* sp1 by Eulin et al. (2017e). Recently *Rhopalodia* was merged with *Epithemia* (Ruck et al. 2016). The new species, *E. boucheziae*, is an example of a new taxon

established using several criteria, following the prerequisites of the integrative taxonomy concept.

Intermediate cases: The second group comprises cases where there is also a relatively easy correspondence between genetic and morphological criteria. However, these cases are more complex because morphologically similar species have been described in the literature, which can lead to misidentifications. Earlier LM observations carried out to establish the regional Atlas of the West Indies (Eulin *et al.* 2017b) identified two morphologically similar species, *G. bourbonense* and *G. designatum*. Careful morphological re-examination of the samples in the present study demonstrated that all the specimens belonged to *G. bourbonense*, as several morphometric features were outside the ranges for *G. designatum*. In addition, the two sequences (ASV11, ASV16) were obtained from the same samples and both were placed in the *G. bourbonense* clade. We therefore concluded that the species in our samples was *G. bourbonense*.

Complex case: The last category concerns groups of several, genetically similar sequences and morphologically similar forms that were impossible to resolve. Three frequent (ASV5, ASV8 and ASV37) and two rarer (ASV57, ASV150) sequences, all belonging to the genus *Nitzschia*, were in the same clade (*N. inconspicua*, *N. amphibia*). Based on the morphology of the specimens present in the samples in which these sequences were frequent, they could match several species (*N. inconspicua*, *N. amphibia*, *N. denticula* and *N. frustulum*), but also morphologically similar forms previously referred to as *N. frustulum* forma 2, *N. frustulum* forma 3, *Nitzschia* sp. 64 and *Nitzschia* sp. 41 (Eulin *et al.* 2017e). In several cases, the read abundance of a single ASV was correlated with several species abundances identified by LM. In addition some ASVs that were not part of the same clade, co-occurred together and were correlated with the same species in LM. There are several possible explanations for this lack of correspondence. For instance, the correspondence between genetic and morphological criteria may sometimes exist only for morphological features which are usually not taken into account for species discrimination (e.g. *Gomphonema parvulum* [Kützing] Kützing in Kermarrec *et al.* 2013). Another explanation is that some morphological species are composed of several cryptic species (and genotypes) that do not co-occur, which hampers the correspondence between morphological and genetic criteria (Trobajo *et al.* 2009). This example clearly shows the limits of Rimet *et al.* (2018) method.

These West Indies islands host more than 100 morphologically identified species. We could establish a clear link with one or several barcodes in Diat.barcode v.10 for seven species, of which four species are illustrated here. Five were simple cases (*U. gouldardii*, *N. boucheziae*, *E. boucheziae*, *Sellaphora nigri* (De

Notaris) C.E. Wetzel & L. Ector, *Navicula incarum* U. Rumrich & Lange-Bertalot), and two were intermediate cases (*G. bourbonense/designatum*, *Navicula escambia/symmetrica/simulata*). These were all essentially abundant species. For rarer species, the method could not be applied. Cell isolation and culturing followed by an integrative taxonomical approach is the only solution to resolving such problematic taxonomic groups.

Conclusion

Metabarcoding is a promising approach that simplifies diatom identification and overcomes the problems associated with the traditional morphological approach (Vasselon *et al.* 2019), however, diatoms exhibit high diversity (e.g. Levkov *et al.* 2007, Mann & Vanormelingen, 2013) and strong endemism (Chonova *et al.* 2021, Verleyen *et al.* 2021, Rimet *et al.* 2023). To correctly identify species and use them effectively for assessing ecological quality, it is necessary to continue exploring their diversity, especially in poorly studied areas, such as the tropics. Moreover, to describe species and accurately define their boundaries, it is necessary to use additional criteria to the morphological criteria, such as molecular criteria (Dayrat 2005).

Work to expand the Diat.barcode library of the West Indies using Rimet *et al.* (2018) methodology now allows the majority of diatom environmental sequences in rivers to be identified. The proportion of sequences identified to species level has been increased from 45% to 84% in the latest version of Diat.barcode (v.10) with all necessary metadata (Rimet *et al.* 2021a, b). Furthermore, our newly described species can easily be identified even in the presence of morphologically sister species (e.g. Evans *et al.* 2009, Rivera *et al.* 2018a, b) since they have associated barcodes. However, even if Rimet *et al.* (2018) method enables quick and cost-effective completion of the reference barcoding library, in some cases it is impossible to apply to a complex taxonomic group, especially when several similar taxa are present in the microscope samples and amplicon data. In this case, we recommend using isolation and culturing methods alongside a careful morphological study.

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Disclosure statement

No potential conflict of interest was reported by the author(s).


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Supplementary data

Supplemental data for this article can be accessed at <https://doi.org/10.1080/0269249X.2023.2237977>. Anne Eulin-Garrigue and Estelle Lefrançois provided the supplemental data when floristic counts were made in the framework of the monitoring of rivers (Eulin et al. 2017a, b, c, d, e, f). Figure legends for Supplementary data: Supplementary data 1. LM micrographs of *Nupela boucheziae* sp. nov. Figs 1–23. Valves with raphe and Figs. 1–19. Valves without raphe, from a site called Blanche-Pont de l'Alma in Martinique. Scale bar = 10 µm. Supplementary data 2. LM micrographs of *Epithemia boucheziae* sp. nov. Figs 1–10. Valve view. Figs 11–13c. Girdle view, from river Riviere Bras David in Guadeloupe. Scale bar = 10 µm. Supplementary data 3. SEM micrographs of *Epithemia boucheziae* sp. nov. Figs 1–2. External valve view. Figs. 3–4 Internal valve view. Scale bars = 10 µm. Supplementary data 4. Spearman's correlation between frustule counts of suspected species and percentage of reads of ASVs. (Correlation coefficient: bottom left of the table, *p*-value: upper right of the table.)

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Figure legends for Supplementary data

Supplementary data 1. LM micrographs of *Nupela boucheziae* sp. nov. Figs 1–23. Valves with raphe and Figs. 1–19. Valves without raphe, from a site called Blanche-Pont de l'Alma in Martinique. Scale bar = 10 µm.

Supplementary data 2. LM micrographs of *Epithemia boucheziae* sp. nov. Figs 1–10. Valve view. Figs 11–13c. Girdle

view, from river Rivière Bras David in Guadeloupe. Scale bar = 10 μm .

Supplementary data 3. SEM micrographs of *Epithemia boucheziae* sp. nov. Figs 1–2. External valve view. Figs. 3–4 Internal valve view. Scale bars = 10 μm .

Supplementary data 4. Spearman's correlation between frustule counts of suspected species and percentage of reads of ASVs. (Correlation coefficient: bottom left of the table, p -value: upper right of the table.)